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Nest site choice by the intertidal spider *Desis formidabilis*
(Araneae: Desidae) and nest utilization by its
hymenopteran egg parasitoid

Running title:

Intertidal spider and parasitoid

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ABSTRACT

1. *Echthrodesis lamorali* Masner 1968 is the only known parasitoid of the eggs of the intertidal rocky shore spider *Desis formidabilis* O.P. Cambridge 1890 and is endemic to a small area of South Africa.
2. The abundance of spider nests and parasitoid presence were assessed in relation to their in- and between-shore location at multiple sites within the distribution of *E. lamorali* along the Cape Peninsula (Western Cape, South Africa).
3. *Desis formidabilis* nests were more abundant in the mid-shore zone than higher or lower up the shore. Spider population sizes also differed between collection sites,

- 25 with higher numbers recorded on the cooler western
26 coast of the peninsula.
- 27 4. Evidence of parasitoid activity was recorded in 43.31%
28 of the 127 nests and 13.85% of the 592 egg sacs they
29 contained.
- 30 5. Where parasitoids gained entry to a spider egg sac,
31 oviposition took place into all of the eggs present.
- 32 6. Incidence of wasp activity was positively correlated with
33 spider nest concentration, not height up the shore,
34 suggesting that the host and parasitoid are tolerant of salt-
35 water inundation.
- 36 7. These results should assist managers of the Table
37 Mountain National Park, in which the full distribution of
38 *E. lamoralis* falls, to better understand this component of
39 rocky shore community dynamics.

40

41 KEYWORDS: Density dependence, *Echthrodesis lamoralis*,
42 Platygastridae, Scelioninae, Rocky shore, Zonation

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INTRODUCTION

48

49 Arthropods have repeatedly colonised intertidal rocky shore
50 environments (Cheng, 1976; Barber, 2009). Crustaceans, a
51 predominantly marine group, are the most conspicuous, but
52 representatives of mainly terrestrial groups such as insects and
53 spiders are also present. Dead and living plant and animal
54 material provides resources for scavengers and predators, which
55 in turn support other invertebrates, including parasitoids
56 (Paetzold et al., 2008; Barber, 2009). Nonetheless, this is a harsh
57 environment to colonise (Lubke, 1998), with each individual
58 shore boasting different characteristics depending on the activity
59 of the tides and currents (Tietz & Robinson, 1974; Lubke, 1998).
60 Single shores alone are also a highly heterogeneous environment
61 from a variety of abiotic aspects (Lubke, 1998). These abiotic
62 factors in turn act as 'stressors' on biotic ones, leading to
63 community structuring (Foster & Treherne, 1976; Sanford,
64 2002).

65

66 Spiders are a generally little-known inhabitant of the intertidal
67 region. A total of eleven spider families have been recorded from
68 intertidal zones globally, with Desis Walckenaer 1937 (Araneae:
69 Desidae) the most common and cosmopolitan genus (McLay &
70 Hayward, 1987). Individuals belonging to this genus also nest
71 lower down the intertidal zone than any other maritime spiders

72 (McLay & Hayward, 1987). *Desis formidabilis* O.P. Cambridge
73 1890 (Araneae: Desidae) (Figure 1) is known only from southern
74 Africa, where it occurs on rocky shores from Lüderitz (Namibia)
75 in the north-west to East London (Eastern Cape, South Africa)
76 in the east (Filmer, 1995; Dippenaar-Schoeman & Jocqué,
77 1997). This species is cribellate, nesting in silk-lined crevices
78 or empty shells found within the intertidal region (Lamoral,
79 1968; McLay & Hayward, 1987) (Figure 1), where it feeds
80 nocturnally on isopods and amphipods (Day, 1974; Dippenaar-
81 Schoeman & Jocqué, 1997, van Noort, 2009).

82

83 The intertidal habitat of *D. formidabilis* may allow the spider to
84 avoid attack from predatory spider-hunting wasps (Pompilidae)
85 (Lamoral, 1968), but in one part of its range, its eggs are host to
86 a specialist egg parasitoid, *Echthrodesis lamorali* Masner 1968
87 (Hymenoptera: Scelioninae) (Lamoral, 1968; Masner, 1968; van
88 Noort, 2009; Owen et al., 2014; van Noort et al., 2014; Owen,
89 2015; Owen et al., 2017) (Figure 2). The distribution of this wasp
90 is restricted to a small area of the Cape Peninsula, south of Cape
91 Town in the Western Cape Province of South Africa (Lamoral,
92 1968; Masner, 1968; van Noort, 2009; Owen et al., 2014; van
93 Noort et al., 2014; Owen, 2015; Owen et al., 2017) and is fully
94 encompassed in the Table Mountain National Park, a
95 conservation region. *Echthrodesis lamorali* is one of only three
96 species of parasitoid wasps worldwide that display unequivocal

97 adaptations to rocky shore intertidal conditions and can be
98 considered as a truly maritime species (van Noort et al., 2014).

99

100 A single *D. formidabilis* nest is built by one female only, and
101 takes the form of several purse-like wedge-shaped egg sacs that
102 fit together within a shell or crevice (van Noort, 2009; van Noort
103 et al., 2014) (Figure 1). Not all egg sacs contain eggs, but where
104 they are laid, one egg mass is produced per sac, and the number
105 of eggs within each varies (van Noort, 2009; van Noort et al.,
106 2014) (Figure 2). The empty egg sacs may act as ‘red herrings’
107 or ‘decoys’, that reduce the searching efficiency of the parasitoid
108 (van Noort, 2009; van Noort et al., 2014). Decoy chambers with
109 this apparent effect have been described in the galls of some
110 species of gall wasps (Hymenoptera: Cynipidae) on *Quercus*
111 spp. (Askew, 1984).

112

113 The intertidal zone represents a challenging and dangerous
114 environment for both the spider and its associated parasitoid.
115 Submersion by seawater is likely to be a particularly important
116 structuring environmental variable and it is hypothesised here
117 that spider nests would be distributed predominantly on the
118 lower shore, which maintains a higher, less variable humidity
119 than upper shore regions, as observed by McLay & Hayward
120 (1987) for the genus. As the parasitoid is of more recent

121 terrestrial origin (van Noort et al., 2014), it is hypothesized that
122 *E. lamorali* would exhibit an opposing in-shore pattern by
123 making use of more spider nests further away from the land-sea
124 interface. This study assesses the factors influencing patterns in
125 the distribution of spider population sizes and parasitism
126 incidence in relation to both their in- and between-shore location
127 at multiple sites within the distribution of *E. lamorali* along the
128 Cape Peninsula (Western Cape, South Africa). Very little is
129 currently known about the ecology of the spider and wasp, and
130 their interactions, despite their unique nature. This study
131 therefore also aimed to describe the within-habitat location and
132 characteristics of the nests of *D. formidabilis* along the Cape
133 Peninsula with emphasis on the nest size and composition, egg
134 numbers and egg success. The full distribution of *E. lamorali* is
135 encompassed in the SANParks Table Mountain National Park.
136 To better manage the park, it is important for conservation
137 officers and managers to understand the dynamics of this unique
138 component of intertidal communities that exists nowhere else.

139

140 MATERIALS AND METHODS

141 Localities

142 Spider nests were collected from four sites located on either side
143 of the Cape Peninsula (Western Cape Province, South Africa)
144 where *E. lamorali* was known to be present: to the west,

145 Kommetjie ($34^{\circ} 8' 22.7034''$ S, $18^{\circ} 19' 17.5794''$ E) and
146 Olifantsbos Point (Table Mountain National Park [TMNP]; 34°
147 $15' 29.6274''$ S, $18^{\circ} 22' 54.0474''$ E) and to the east, Buffels Bay
148 (TMNP; $34^{\circ} 19' 19.5594''$ S, $18^{\circ} 27' 44.028''$ E) and Simon's
149 Town ($34^{\circ} 9' 43.7394''$ S, $18^{\circ} 25' 55.5234''$ E) (Owen et al.,
150 2014; Owen, 2015) (Figure 3).

151

152 The rocky shore at each site was separated into three zones: the
153 sublittoral fringe, eulittoral range and littoral fringe (zones A, B
154 and C respectively), following Lamoral (1968). Zones were
155 demarcated using visual assessment of the coastline during the
156 changing tides, in combination with patterns in encrusting
157 materials on the shore, such as algal clusters.

158

159 Nest collection

160 Nests were collected on six occasions at the lowest tides every
161 two weeks in spring between 9 September and 22 November
162 2014. A single transect perpendicular to the shoreline was placed
163 at random along the shoreline during each collection. The
164 transects ran from the upper littoral (Zone C) to the lower
165 sublittoral fringe (Zone A), encompassing all of the rocky shore
166 in-between, and thus ranged in length depending on the shore
167 sampled, the physical aspects of which were highly variable

168 between sites due to the heterogeneous nature of the intertidal
169 region. The area within a width of 2m of the line within each of
170 the three zones was then visually searched for 30 minutes. Each
171 collected *D. formidabilis* nest was stored individually in mesh-
172 covered tubes and all nest characteristics, described below, were
173 recorded.

174

175 Nest contents

176 In the laboratory, each nest was separated into its constituent egg
177 sacs, and the number, condition and contents of each were
178 recorded. Unhatched eggs were left within their individual egg
179 sacs (to maintain adequate humidity levels for the eggs) in
180 individual glass rearing chambers, the openings of which were
181 sealed with netting. These chambers were kept at room
182 temperature and moistened daily by adding a few drops of
183 seawater to assist with the maintenance of humidity. The eggs
184 were given three months to incubate and allow for the
185 development and emergence of embryos therein. The state of the
186 eggs at the time of collection were therefore referred to as ‘pre-
187 incubation’, while those that had been maintained in the
188 laboratory for an incubation period were thereafter referred to as
189 ‘post-incubation’ eggs. Wasps and spiders that emerged from the
190 eggs were counted and sexed (in the case of any *E. lamorali*
191 emergence), while the contents of any eggs that did not hatch

192 were subsequently assessed under a binocular dissecting
193 microscope in an attempt to determine the species of the
194 unsuccessful embryo where possible. Eggs that gave rise to
195 wasps, or in which failed wasp embryos were identifiable were
196 classed as being ‘parasitized’. All material was stored in 96%
197 ethanol and accessioned into the entomology wet collection at
198 the Iziko South African Museum (Cape Town, Western Cape,
199 South Africa).

200

201 Statistical analyses

202 To explore the factors influencing patterns in the distribution of
203 spider population sizes and parasitism incidence, they were
204 assessed using two different sample size-corrected, Akaike
205 Information Criterion (AICc, which are better suited to small
206 datasets, such as those used here, than AICs) based, reverse
207 model building processes (Symonds & Moussalli, 2011) in R
208 version 3.1.3 (R Core Team, 2015) using the packages MuMIn
209 (Barton, 2015) and car (Fox & Weisberg, 2011). An AICc
210 analysis was chosen as it allows exploration of the system as a
211 whole, and indicates what factors interact to define the
212 ecological patterns that are evident. The most complex model
213 using all possible predictor (independent) variables (those
214 recorded for each nest during sample collection) was first
215 constructed and co-linearity within the variables was determined

216 for each outcome (spider population size or parasitism
217 incidence). Where co-linearity was detected, the variable
218 indicated (GVIF>5) was removed. Akaike Information Criterion
219 values (AICcs) were then calculated for all possible models
220 using the variables from the initial one. All models developed in
221 this process with a delta value of less than 2 were considered and
222 the variables used in the construction of these was assessed as in
223 Symonds & Moussalli (2011). The most commonly occurring
224 predictors that made the most biological sense were used in
225 construction of the final model that outlined the effect that the
226 remaining predictor variables had on the spider population size
227 or parasitism incidence (Zuur et al., 2010; Symonds &
228 Moussalli, 2011).

229

230 To answer specific questions on areas of interest, the following
231 statistical tests were completed in Statistica 10 (StatSoft Inc.,
232 2011). Differences in the number of egg sacs and the number of
233 eggs laid per nest and egg sac collected from different sites and
234 zones were analysed using a Kruskal Wallis test. Regression
235 models tested for correlations between the total number of egg
236 sacs and how many were left unused, and between the number
237 of eggs and male wasp presence in the post-incubation emerged
238 wasps. Generalized linear models revealed whether differences
239 existed between sites and zones for hatching success, which is
240 defined here as the full development and post-incubation

241 hatching of either a spider or wasp from an egg, of parasitized
242 and non-parasitized eggs.

243

244 RESULTS

245 All AICc results tables are presented as supplementary
246 information in a supporting document. Only the patterns and
247 interpretation of these results are presented here.

248 Nest numbers

249 A total of 127 nests comprising 592 egg sacs were collected from
250 the four sites. The AICc model for spider population size (the
251 number of nests collected) returned a total of eight plausible
252 models with deltas of less than two. The final model illustrated
253 that spider population sizes differed significantly between zones
254 and sites. The most nests were collected in zone B, followed by
255 A, and finally zone C, although zones A and B did not differ
256 significantly (Figure 4A). The sites on the western shore of the
257 Peninsula hosted the most spiders, with the highest population
258 density (most number of nests collected) at Kommetjie, then
259 Olifantsbos, followed by Buffels Bay and finally Simon's Town
260 (Figure 4B).

261

262 Nest contents

263 The number of egg sacs per nest ranged from one to 23 (mean \pm
264 SE = 4.0 ± 0.34), and differed between sites ($H_{(3, 127)} = 10.430$,
265 $P = 0.015$), but not zones ($H_{(3, 127)} = 2.107$, $P = 0.349$). Larger
266 nests were found at Olifantsbos than Kommetjie, but no
267 differences existed between the other sites (Figure 5). Of the 127
268 nests collected, 28 (22.05%) consisted of only a single egg sac.

269

270 Among the 99 collected nests that had two or more egg sacs, 12
271 (12.12%) contained no eggs in any of the egg sacs, most
272 (51.52%) had a mixture of used (with spider eggs) and unused
273 (no contents) egg sacs, and 36 (36.36%) nests were collected
274 where all egg sacs were used. Overall, in all of the collected
275 nests, 42% of the egg sacs were left unused (Figure 6), 9% of
276 which showed evidence of parasitoid action (holes chewed by
277 wasps through the silk) (Figure 6A). The relationship between
278 the total number of egg sacs in a nest and the number where no
279 eggs had been laid was positively correlated ($F_{(1,125)} = 262.61$, P
280 < 0.0001 , $R^2 = 0.678$), indicating that bigger nests contained
281 more unused egg sacs.

282

283 Thirty-three percent of pre-incubation egg sacs contained one or
284 more eggs (Figure 6), of which 67% were still viable (“fresh” or
285 “brown unhatched”) and were isolated for incubation (Figure
286 6B). The mean number of eggs per nest was 41.11 ± 4.439 (SE)

287 (minimum = 1, maximum = 226), while the mean in egg sacs
288 where at least one egg had been laid was 19.15 ± 0.803 (SE),
289 with a minimum of one egg and a maximum of 54. The number
290 of eggs per egg sac differed significantly between sites ($H_{(3, 85)} =$
291 13.455 , $P = 0.004$), but not zones ($H_{(2, 85)} = 1.218$, $P = 0.544$),
292 with spiders at Olifantsbos producing more eggs than at Buffels
293 Bay (Figure 7). The remaining used egg sacs all contained
294 spiderlings (21%) or wasps (4%) (Figure 6). Most of the former
295 were alive (98%), with 60% darker and older, and 38% freshly
296 hatched (Figure 6C), while the majority (88%) of the latter where
297 the egg sacs contained wasps hosted live adult wasps (Figure
298 6D).

299

300 Parasitism of eggs, i.e. the number of eggs per egg sac from
301 which wasps emerged or were visible as failed embryos, in a
302 single egg sac ranged from 11.36% to 100%, with a mean of 60%
303 ± 5.45 (SE). The parasitism incidence AICc returned zone as the
304 only significant predictor as to whether an egg sac was
305 parasitized or not. Following a similar trend to the spider
306 population size modelled previously, most parasitism took place
307 in the eulittoral zone (B), followed by the sublittoral fringe (A)
308 and lastly the littoral fringe (C) (Figure 8).

309

310 Post-incubation figures and sex ratios of resultant wasps

311 Isolated non-parasitized eggs exhibited a $47.99\% \pm 10.40$ (SE)
312 hatching success rate in the laboratory. Parasitized eggs ($n=704$
313 eggs) were less successful, with an average hatching rate of
314 $24.13\% \pm 8.61$ (SE). In most cases where parasitized eggs failed,
315 the wasp embryo reached full development, but never eclosed.
316 No significant differences were detected between either of these
317 success rates at all sites (not parasitized: $F_{(3, 62)} = 0.291$, $P =$
318 0.832 ; parasitized $F_{(2, 45)} = 0.444$, $P = 0.644$) and zones (not
319 parasitized $F_{(2, 62)} = 2.108$, $P = 0.130$; parasitized $F_{(2, 45)} = 1.138$,
320 $P = 0.330$).

321

322 A total of 83 wasps were reared from the collected nests, 67
323 (80.72%) of which were female and only 16 (19.28%) male,
324 indicating a strong female bias. A mean of 0.89 ± 0.29 (SE) male
325 wasps were found per egg sac, while females averaged $3.72 \pm$
326 1.10 (SE) individuals per egg sac. The number of eggs in an egg
327 sac and the number of males that emerged from them were
328 marginally correlated ($R^2_{(16)} = 0.211$, $P = 0.055$), with a trend for
329 larger broods to contain more male wasps.

330

331 DISCUSSION

332 This study has shown that *D. formidabilis* favours the western
333 coastline of the Cape Peninsula, which is known to be much

334 cooler than the east (Teske et al., 2006; Teske et al., 2011). It
335 also favours the mid-shore zones, where twice a day salt-water
336 inundation occurs, but for a shorter duration than below it, and
337 longer duration than above it. Nests of the spiders varied greatly
338 in terms of the numbers of egg sacs and eggs that they contained.
339 Many egg sacs were empty and hatching success rates among the
340 eggs was low. Within individual sites, parasitism of the eggs by
341 *E. lamoralis* followed a density-dependent pattern, with a
342 concentration in the mid-shore zone where most hosts were
343 located. The wasp population sex ratios were highly female-
344 biased.

345

346 Significant differences in spider population sizes existed
347 between intertidal zones and sites. In contrast to the original
348 hypothesis that the spiders would prefer the lower regions of the
349 shore closer to the water's low tide edge, the intermediate
350 eulittoral zone and highest sublittoral fringe hosted the largest
351 spider populations and the fewest spider nests were found lowest
352 down in the littoral fringe. This distribution pattern is likely to
353 reflect the influence exerted by environmental conditions on a
354 shore. Conditions become drier and hotter for longer periods of
355 time up the intertidal region towards the terrestrial environment
356 due to the movements of the tides (Sanford, 2002). Lower down
357 the intertidal region, however, the physical force of wave action
358 can be too stressful for certain species (Sanford, 2002). These

359 abiotic changes across the intertidal range have large effects on
360 the in-shore distributions of most of the organisms living in this
361 environment, and this appears to include the spatial distribution
362 of *D. formidabilis*.

363

364 The populations of the spider on the west coast are also
365 significantly larger than those on the eastern side of the
366 peninsula. A large variety of environmental and community-
367 based parameters can have an effect on population sizes (Foster
368 & Treherne, 1976; Sanford, 2002). This is often striking for
369 intertidal regions, where the combination of varying biotic and
370 abiotic conditions on different shores can lead to unique patterns
371 (Tietz & Robinson, 1974; Lubke, 1998). As discussed in Owen
372 et al. (2014) and Owen (2015), the spider populations fall within
373 the ecotone between the warm temperate biogeographical region
374 to the east and the cool temperate region to the west. The
375 differences between these two biogeographical zones have
376 strong influences on local conditions, and consequently modify
377 community composition on the two shores (Tietz & Robinson,
378 1974; Foster & Treherne, 1976; Lubke, 1998; Sanford, 2002).

379

380 A significant outcome of the model on spider population sizes is
381 that parasitism was not identified as a controlling factor. In many
382 other systems, parasitoids exert a top-down controlling effect on

383 spider populations, such as in spider populations affected by
384 pompilid wasp abundances on islands in the Gulf of California
385 (Polis & Strong, 1996), and the web building spiders attacked by
386 *Trypoxylon figulus* von Linné 1758 (Hymenoptera:
387 Crabronidae) in Switzerland (Coudrain et al., 2013). In contrast,
388 the harsh environmental conditions and steep environmental
389 gradients of the intertidal habitat of *D. formidabilis* result in
390 dominance by bottom-up dynamics, which incorporate both
391 biotic and abiotic factors (Chen & Wise, 1999; Denno et al.,
392 2002; Menge et al., 2015). A bottom-up controlled system is
393 defined as one where food sources and environmental conditions
394 have a larger effect on a population than the effect of predators
395 and parasitoids (Lynam et al., 2017). Such bottom-up control has
396 previously been recorded in detritus-driven ecosystems such as
397 the intertidal zone (Chen & Wise, 1999; Menge et al., 2015) and
398 also among other arachnid species living in North American
399 intertidal marshes (Denno et al., 2002).

400

401 Although differences existed in the average nest sizes (number
402 of egg sacs) between some sites in this study, no clear difference
403 between eastern and western shores of the Peninsula was
404 discernible. However, the number of eggs produced per nest on
405 the west coast of the Peninsula was higher than in the east.

406

407 The average *D. formidabilis* nest comprised around four egg
408 sacs, which is comparable to the nests produced elsewhere by
409 another maritime species in the same genus, *Desis marina*
410 Hector 1877 (Araneae: Desidae), which contained an average of
411 3.4 egg sacs (McLay & Hayward, 1987). Almost half of all egg
412 sacs in the *D. formidabilis* nests were unused. This contrasts with
413 the results of McLay & Hawyard (1987), who recorded no
414 unused egg sacs in the nests of *D. marina*. However, the number
415 of eggs in utilised egg sacs were similar between the species,
416 with the number of eggs in a single *D. marina* egg sac ranging
417 from 11-61, compared with 1-54 eggs in *Desis formidabilis*.

418

419 Nine percent of the unused egg sacs exhibited signs of *E.*
420 *lamorali* activity (chewed holes). The “red herring” hypothesis
421 postulated by van Noort (2009) and van Noort et al. (2014)
422 suggested that wasps may not have enough energy resources left
423 to enter a second egg sac after having chewed into a first that
424 contains no eggs. Austin (1984) showed that *Ceratobaeus*
425 *masneri* Austin 1983 (Hymenoptera: Platygasteridae) females
426 usually die shortly after producing their first brood because dead
427 individuals were often found on or in the host’s nests, though
428 this may be the result of starvation during the brood guarding
429 that is displayed by some platygasterids following oviposition
430 (Matsuo et al., 2014). In contrast, none of the empty *D.*
431 *formidabilis* egg sacs that exhibited signs of parasitoid entry

432 contained dead adult wasps, which casts doubt on the ‘red
433 herring’ theory as originally postulated, although any time
434 ‘wasted’ by searching females is likely to reduce their lifetime
435 reproductive success.

436

437 Parasitism incidences within a single egg sac averaged at 60%,
438 in contrast to van Noort et al. (2014) on the same two species,
439 who consistently recorded a parasitism incidence of 100% within
440 a single egg sac. However, of the 566 egg sacs assessed in this
441 study, only one contained both spiderlings and wasps and given
442 that encapsulated and putrefied embryos were quite frequent,
443 initial parasitism rates may often be close to 100% within those
444 host egg batches utilised by the wasp. This parasitism frequency
445 is not common for the Scelioninae (van Noort et al., 2014).

446

447 The zone in which a nest was built was the only factor identified
448 as significantly influencing parasitism rates, with most
449 parasitism recorded in the middle zone, the lowest in the upper
450 zone and the lowest zone intermediate between the others.
451 Parasitism incidence is likely to be affected by the wasp’s ability
452 to find and recognise the host nest, a behaviour that is influenced
453 by a variety of environmental factors (Fox et al., 1990; Austin et
454 al., 2005). In this case, in contrast to the original hypothesis, the

455 parasitism incidence followed density patterns displayed by the
456 host spider rather than abiotic factors like water inundation.

457

458 For *D. marina* eggs, McLay & Hayward (1987) reported a high
459 hatching success rate of 89.99%. In this study, most sites
460 experienced an approximate 50% success rate of juvenile spider
461 emergence from eggs that had not been parasitized. A low
462 hatching success rate of eggs is common for species with high
463 reproductive outputs (Deevey, Jr., 1947; Murdoch, 1966; Gadgil
464 & Bossert, 1970), such as *D. formidabilis*.

465

466 *Echthrodesis lamorali* have strongly female-biased broods (83%
467 female) in this study, similar to the 87.29% female brood
468 recorded by van Noort et al. (2014). Female dominated sex ratios
469 are common for platygastriids (Austin, 1984; Strand, 1988; Fox
470 et al., 1990; Austin et al., 2005). As overall clutch size increases,
471 Green et al. (1982) showed that the number of males follows the
472 same trend for other parasitoid wasps, which was also
473 demonstrated here, although only weakly.

474

475 In conclusion, *D. formidabilis* displays a preference for mid-
476 shore elevations along cooler rocky coastlines. Its egg parasitoid,
477 which has a very restricted distribution within the wider range of

478 its host, is able to utilise spider nests from the lower to upper
479 shores, but concentrates its activities in the mid-shore, where its
480 hosts are most abundant. These results should assist managers of
481 the Table Mountain National Park in better understanding the
482 intertidal community dynamics within the reserve.

483

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497

498 CONTRIBUTIONS OF AUTHORS

499 Candice Owen, Julie Coetzee and Simon van Noort were
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501 of the fieldwork, data collection and manuscript preparation.
502 Stephen Compton assisted with manuscript content and data
503 analysis.

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FIGURE LEGENDS

660 Figure 1: *Desis formidabilis* lives solely in the intertidal habitat.

661 (A) The shore at Kommetjie where *Echthrodesis lamorali* was

662 first discovered. The different rock colours visible in this

663 photograph indicate the three zones. (B) The typical habitat in

664 which both *D. formidabilis* and *E. lamorali* live. (C and D) *Desis*

665 *formidabilis* constructs its nests in gaps between rocks and under

666 empty limpet shells, which are usually concealed beneath the

667 boulders shown in the previous photo, using silk to form a

668 waterproof lining between the substrate and nest. (E) *Desis*

669 *formidabilis* individuals within a nest. Pale pink egg sacs are

670 visible to the bottom and right of the nest. (F) A close-up

671 photograph of the egg sacs showing the purse-like shape that

672 opens when spiderlings exit the egg sac. The entry or exit holes

673 made by *E. lamorali* are visible on the two left-most egg sacs.

674 Figure 2: (A) *Desis formidabilis* eggs within two opened egg

675 sacs showing freshly laid white eggs and mature parasitized

676 brown eggs. Egg colour was shown to correlate with parasitism

677 status, with the brown colour of the eggs on the left a good

678 indicator of parasitism. (B) Freshly hatched *D. formidabilis*
679 spiderlings are pale and emerge through the opened edge of the
680 egg sac. (C and D) *Echthrodesis lamorali* oviposits one egg into
681 each *D. formidabilis* egg in an egg sac. (E) *Echthrodesis*
682 *lamorali* males eclose first and guard an egg containing a female.
683 On her emergence, the two immediately mate. (F) The full
684 habitus of an *E. lamorali* female.

685 Figure 3: The four sites that were actively searched for *Desis*
686 *formidabilis* and *Echthrodesis lamorali* presence, Kommetjie
687 (A), Olifantsbos Point (B), Buffels Bay (C) and Simon's Town
688 (D).

689 Figure 4: The mean number (\pm SE) of *Desis formidabilis* nests
690 collected from different zones (A) and sites (B) throughout all of
691 the collection trips (letters indicate significant differences).

692 Figure 5: The mean number (\pm SE) of egg sacs recorded in *Desis*
693 *formidabilis* nests collected from different sites (letters indicate
694 significant differences).

695 Figure 6: The pre-incubation contents of 592 collected *Desis*
696 *formidabilis* egg sacs. Egg sacs with no contents (A) were either
697 intact, or had holes chewed by adult *Echthrodesis lamorali*. Eggs
698 (B) found in egg sacs at the time of collection exhibited the
699 greatest variety of states, with the majority comprising of freshly
700 laid, live eggs, followed by older, brown unhatched eggs,
701 batches inflicted with mould, hatched eggs and finally putrefied

702 eggs. Spiders (C) present in egg sacs at the time of collection
703 were classed as either live, freshly hatched individuals; live,
704 older spiderlings; or mouldy dead adults. Where egg sacs
705 contained wasps at the time of collection (D), they were either
706 dead or alive.

707 Figure 7: The mean number (\pm SE) of eggs in egg sacs of *Desis*
708 *formidabilis* nests collected from different sites (letters indicate
709 significant differences).

710 Figure 8: Percentage parasitism of *Desis formidabilis* eggs in
711 different egg sacs by *Echthrodesis lamorali* differed
712 significantly between zones (letters indicate significant
713 differences).















